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Flexible percolation fibrous thermal insulating composite membranes for thermal management†

Lu An,^a Massimiliano Di Luigi,^a Jingye Tan,^a Danial Faghihi^a and
Shenqiang Ren^{b,c}

Flexible thermal insulating membranes are ubiquitous in thermal management. Nevertheless, difficulties arise for composite membranes to combine a resilient, robust structural framework with uniform percolation networks purposefully conceived for thermal insulation. Herein, by controlling the microstructure homogeneity, we report flexible, hydrophobic thermal insulating membranes consisting of ceramic fiber and porous silica materials. The resulting nanofibrous membrane composites exhibit a low thermal insulation of $11.4 \text{ mW m}^{-1} \text{ K}^{-1}$, a low density of 0.245 g cm^{-3} , mechanical flexibility with a bending rigidity of 1.25 cN mm^{-1} , and hydrophobicity with a water contact angle of 144° . These nanofibrous-reinforced, silica-aerogel-based nanocomposite membranes are potential candidates for advanced thermal management applications.

1 Introduction

Thermal insulating materials are essential for thermal management in buildings, electronics, and wearable systems.^{1–4} The thermal insulating performance of these materials originates from the percolation porous networks, *i.e.* silica aerogels, with narrow pore size distribution, efficiently limiting the gaseous thermal conduction.^{5–8} However, silica aerogels show inherent weak interparticle connections causing poor mechanical performance. The incorporation of fibers could improve porous materials' mechanical performance, while the thermal insulating performance could be degraded by the non-uniform microstructures.^{9–15} The occurrence of an accelerated heat transfer through fibers and large void spaces, as part of solid phase thermal conduction, dramatically limits the thermal insulating performance of fibrous composite materials and their subsequent structural instability under applied mechanical stress.^{16,17}

In this work, we report flexible, hydrophobic thermal insulating membrane nanocomposites consisting of ceramic fibers and silica aerogels. Through a meticulous control of the microstructure of silica aerogels and fiber networks, these thermally insulating membrane nanocomposites achieve mechanical robustness and flexibility. Highly thermal insulating silica aerogels interfacing with fibers effectively restrain the heat transfer from fiber to fiber, which provide a path to preserve the high thermal insulating properties of the composite material.^{4,18–20} Hydrophobic surface modification is carried out by using 1H, 1H, 2H, 2H-perfluorodecyltrichlorosilane (PFDTs). The resulting nanofibrous membrane composites exhibit a low thermal insulation of $11.4 \text{ mW m}^{-1} \text{ K}^{-1}$, a low density of 0.245 g cm^{-3} , mechanical flexibility with a bending rigidity of 1.25 cN mm^{-1} , and hydrophobicity with a water contact angle of 144° . These nanofibrous-reinforced, silica-aerogel-based nanocomposite membranes are promising alternatives in applications related to advanced thermal management.

^a Department of Mechanical and Aerospace Engineering, University at Buffalo, The State University of New York, Buffalo, New York 14260, USA.

E-mail: shenren@buffalo.edu

^b Department of Chemistry, University at Buffalo, The State University of New York, Buffalo, NY 14260, USA

^c Research and Education in Energy, Environment & Water (RENEW), University at Buffalo, The State University of New York, Buffalo, NY 14260, USA

† Electronic supplementary information (ESI) available: Details of the microstructure (including TEM images), surface area, pore width, average particle size, structural characterization (XRD spectra), and the thermogravimetric study of aerogel and composite specimens, respectively; in addition to assessment of hydrophobic performance, SEM images, thermal conductivity, and mechanical properties of the composite membrane. See DOI: <https://doi.org/10.1039/d2ma00915c>

2 Materials and methods

2.1 Fibrous nanocomposite membranes

The whole process includes (1) mesoporous silica powder preparation, (2) ceramic fiber treatment, and (3) fibrous nanocomposite membrane fabrication.

The synthesis of tetraethyl orthosilicate (TEOS) based mesoporous silica powders used in this study follows a previous work.¹⁰ The following materials have been used during the synthesis of silica powders: tetraethyl orthosilicate (TEOS-[Si(OC₂H₅)₄]; ≥99.0%



(GC) [Sigma Aldrich]); cetrimonium bromide (CTAB – [C₁₉H₄₂BrN; high purity] VWR)]; 3,3',5,5'-tetramethylbenzidine (TMB- [–C₆H₂(CH₃)₂-4-NH₂]₂; ≥99%] Sigma Aldrich). The chemical molar ratio of TEOS:CTAB:Urea:TMB:H₂O is 1:0.122:2.21:0.063:49.25, respectively. The sol-gel process with an initial pH of 8 is carried out below 60 °C. Urea could decompose into ammonia for promoting the pH during the gelation process, and carbon dioxide resists the pore collapse. The ambient pressure dried samples would be sintered at 600 °C for 1 h to remove the chemical residue. The powders would be blended in water for 15 seconds and then treated with high frequency ultrasound for less than 30 min. The original ceramic fibers are dispersed in water using a homogenizer and blended using a commercial blender for 15 seconds to make the fibers short. These fibers would be washed with diluted HCl solution (1% v/v).

The solutions of the above powders and fibers are mixed using a homogenizer and under high frequency ultrasound. This ultrasound process would help in the deposition of porous silica onto the acid treated fiber surface. The composite membrane is fabricated through vacuum filtration, and the wet membrane covered by the PET film is loaded onto a hot press stage at 60 °C and 0.3 MPa pressure for 3 hours, which is performed to densify and flatten the membrane. Afterwards the composite membrane covered by two rigid paper and a metal plate is transferred to the oven for drying to resist the buckling. The porous silica mass concentration is adjusted by the mass weight of the used porous silica powders and fibers mentioned above. The thickness is adjusted by the amount of total mixture solution through the control of the porous silica concentration ratio. The hydrophobic surface modification is carried out by depositing 1H,1H,2H,2H-perfluorodecyltrichlorosilane (PFDTs) onto the membrane composites. The sintering treatments at different temperatures (*i.e.*, 150 °C, 200 °C, and 400 °C) are compared with the dried membrane samples.

2.2 Microstructural characterization

A Carl Zeiss AURIGA scanning electron microscope (SEM) and a JEOL 2010 high-resolution transmission electron microscope (HRTEM) are used to characterize the microstructures of the porous silica and composite membranes. The X-ray diffraction patterns of samples were obtained using a Rigaku Ultima IV with an operational X-ray tube power of 1.76 kW (40 kV, 44 mA) with Cu K α radiation ($\lambda = 1.54184 \text{ \AA}$). BET analysis is performed on a Tristar II 3020 (Micromeritics Corp. Atlanta, GA). The specific surface area (SSA) and the pore size distributions are evaluated with the low-temperature nitrogen adsorption-desorption isotherm measurement method. The pure aerogels are degassed at 200 °C for 2 hours before analysis. The surface areas are calculated with the Brunauer–Emmett–Teller (BET) theory using the isotherm adsorption data at P/P_0 from 0.05 to 0.30. The pore size is calculated from the adsorption branch based on the Barrett, Joyner, and Halenda (BJH) method with the average pore diameter calculated using equation $d = 4 V A^{-1}$.

2.3 Thermal performance measurement

The thermal conductivity of the composite membranes is determined using a Thermtest HFM-100 following the ASTM

C518 standard. Pertinent calibration is performed on commercial extruded polystyrene boards of an appropriate thickness and with a known value of thermal conductivity. The membrane thickness is automatically determined using the HFM-100. The measurement begins by fixing the upper and lower plates at 30 °C and 40 °C, respectively, and the thermal conductivity is determined when the heat flux becomes a constant value. Infrared images are recorded using a Fotric 225 Pro Thermal Camera.

2.4 Mechanical performance measurement

The flexibility of membranes with different thicknesses is evaluated using the vertical ring bend method and specimens with dimensions of 50 mm \times 50 mm are evaluated with an ultra-sensitive load cell on a universal test frame (Model DSTM-50KN from United Testing Systems).

3 Results and discussion

The hierarchical fibrous structure combined with silica aerogels ensures superlative thermal insulating performance, while at the same time achieving structural integrity and mechanical flexibility (Fig. 1a). The TEM images show the hierarchical mesoporous silica aerogel and the aerogel/fiber interface, respectively. The silica aerogel materials are shown in Fig. S1 (ESI[†]), which exhibit the existence of a homogeneous mesoporous silica network that effectively percolates through the voids between the structural fiber network originating from the ceramic fibers, thus giving rise to a uniform fiber/silica aerogel framework. This homogeneous microstructure circumvents the heat transfer from the layered structure^{21,22} while the uniformly distributed silica aerogel provides an effective thermal barrier to potential heat transfer among fibers.^{19,22} The aforementioned fiber network mostly comprises short fibers with an average length of around 100 μm as highlighted in Fig. 1f (cross-section of a membrane specimen), which promotes a much larger uniformity of the fiber/aerogel structural network than the one that could be achieved with long threads of fiber due to a greater intricacy between short fibers and mesoporous silica aerogel particles. Moreover, a noticeable flexibility is achieved in the composite membrane by virtue of the resulting entanglement between fibers and porous silica, as highlighted in Fig. 1b.

On the other hand, inducing resistance to moisture on the fibrous membrane could improve its thermal insulating performance under humid conditions, which is achieved by means of hydrophobic surface modification through PFDTs treatment that results in a water contact angle of 144° (inset of Fig. 1b). The in-plane and cross-sectional SEM images (Fig. 1e and f, respectively) show the membrane microstructure featuring an extremely uniform distribution of the non-layered silica aerogel and fiber network structure. Furthermore, the presence of porous fibrous networks significantly reduces the solid phase thermal conductivity within the membrane, hence enhancing its thermal insulating performance.



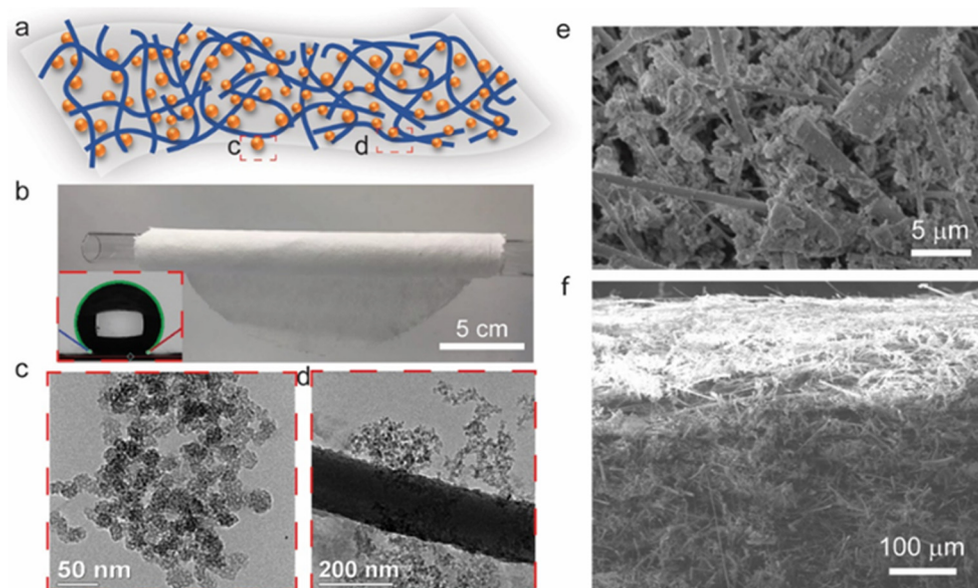


Fig. 1 (a) Scheme of the fiber–aerogel composite membrane. The insets c and d show the corresponding TEM images of mesoporous silica aerogels (c), and fiber-aerogels (d). (b) Flexibility demonstration of the membrane. The inset shows the water contact angle of 144° after hydrophobic coating. (e) In-plane SEM image of the fiber-aerogel composite membrane. (f) Cross-sectional SEM image of the membrane.

Thermal insulation of nanocomposite membrane is derived from the restrained solid thermal conduction through fibers, as well as through effective restriction of the gaseous heat transfer among the matrix. The fibrous percolation networks provide a platform to reduce the solid heat transfer, while silica aerogel coating could block the heat transfer from fiber to fiber. Such a hierarchical architecture obtained in this type of composite materials becomes critical during the improvement of their thermal insulating performance. The sought-after homogeneous network can then be achieved in membranes with the thickness between $500\ \mu\text{m}$ and $1.3\ \text{mm}$ (Fig. 2a and b), with a thickness dependence developing for materials beyond this range. The microstructure of the thickness-dependent composite materials is easily affected by the filtration process, where the layer-by-layer deposition occurs along the fluid flow and gravity direction. Larger gaps and voids between layers could increase the gaseous heat transfer that translates into a reduced thermal insulating performance. The thermal conductivity of membrane nanocomposites with thickness in the region of $300\ \mu\text{m}$ falls within $10\text{--}15\ \text{mW m}^{-1}\ \text{K}^{-1}$, showing a roughly linear increment as the thickness increases up to $1.0\ \text{mm}$. Subsequently, the thermal conductivity increases to $23\text{--}27\ \text{mW m}^{-1}\ \text{K}^{-1}$ as the thickness increases above $1.0\ \text{mm}$ (Fig. 2c).

The incorporation of silica aerogels serves as the thermal barrier within the fibrous percolation networks. Therefore, the thermal conductivity of nanocomposite membranes with similar thicknesses would decrease as the concentration of silica aerogel increases (Fig. 2c and d). The membranes show $\sim 33\%$ lower thermal conductivity when the concentration of silica aerogels increases from $40\ \text{wt\%}$ to $80\ \text{wt\%}$. Even though a higher loading of silica aerogels benefits thermal insulating

performance, the structural integrity of membranes needs to be considered, especially for the membrane with $80\ \text{wt\%}$ silica aerogels. The mass density of the nanocomposite membranes is strictly related to the unit volume of silica aerogels and fibrous networks, with a clear trend showing a decrease in thermal conductivity as the density increases (Fig. 2e). This is probably induced by the gaseous phase component through the fiber/silica aerogel structural network, where a membrane with lower density may feature a higher gaseous phase component of thermal conductivity.

The fibrous nanocomposite membranes with different concentrations of silica aerogels ($33\ \text{wt\%}$, $50\ \text{wt\%}$ and $67\ \text{wt\%}$) and thicknesses ($300\ \mu\text{m}$, $600\ \mu\text{m}$, and $800\ \mu\text{m}$) show great potential for heat resistance (Fig. 3a). Infrared images in Fig. 3b–d are recorded with the surface hot plate kept at different temperatures (*i.e.*: $35\ ^\circ\text{C}$, $47\ ^\circ\text{C}$, and $52\ ^\circ\text{C}$). As the thickness of the nanocomposites increases, the temperature of their top surface decreases, while increasing the concentration of silica aerogels induces a further reduction of the top surface temperature (Fig. 3b–d). Fig. 3e summarizes the temperature difference (ΔT) between the top surface of the fibrous nanocomposite membrane and that of the surface hot plate. Under dissimilar heating temperatures, membranes with different thicknesses have a noticeable temperature difference ΔT , such as $1.5\ ^\circ\text{C}$ for $33\ \text{wt\%}$ with $300\ \mu\text{m}$ thickness compared with $8.5\ ^\circ\text{C}$ for $33\ \text{wt\%}$ with $800\ \mu\text{m}$ thickness. On the other hand, for membranes with similar thicknesses, the temperature difference ΔT is larger at higher temperatures, while higher concentrations of silica aerogels show a larger ΔT , especially for the thickness of $300\ \mu\text{m}$ and $600\ \mu\text{m}$.

The mechanical flexibility of thermal-insulating composite membranes can be affected under different external load



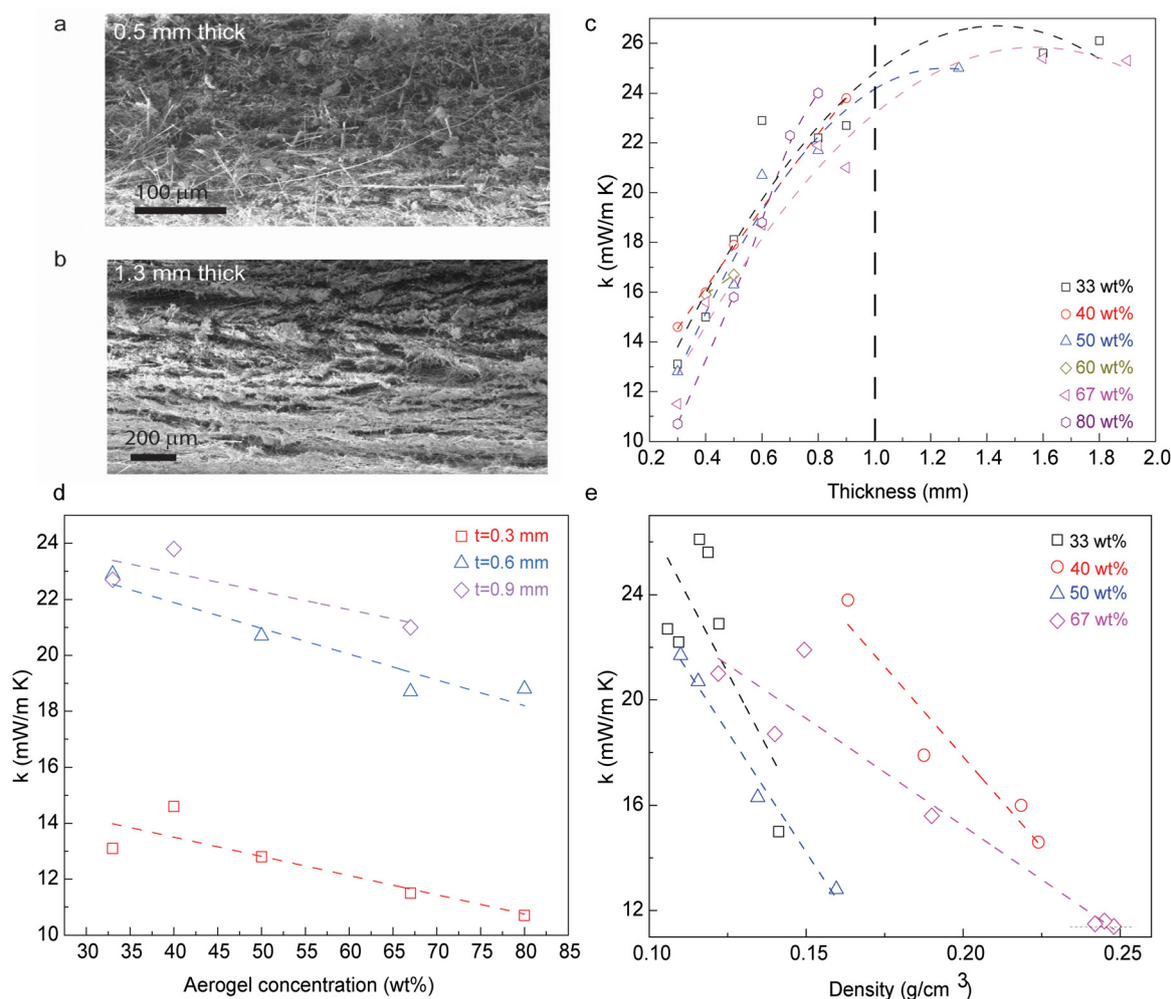


Fig. 2 (a) Cross-sectional SEM image of the membrane with a thickness of 500 μm . (b) Cross-sectional SEM image of the membrane with a thickness of 1.3 mm. (c) Thermal conductivity vs. thickness of the membrane with different concentrations. (d) Thermal conductivity vs. aerogel concentration for membranes with thicknesses of 0.3 mm, 0.6 mm and 0.9 mm. (e) Thermal conductivity vs. density for membranes with different aerogel concentrations.

conditions. The fiber entanglement is a key aspect for preserving the integrity of the structure network bearing loads/stresses, while the type of microstructure of the membrane would also affect its mechanical performance. Here, the mechanical flexibility of the nanocomposite membranes is evaluated using a multi-cycle bend method, which is shown in Fig. 4. After the top plate comes into contact with the membrane's surface, it compresses the shell-like membrane ring (vertical height of 15 mm) from 'point contact' to 'flat-surface contact' with the increase of both displacement and load force simultaneously (Fig. 4a). During the unloading process, the bent membrane would recover back within the elastic region. For a 600 μm thick membrane with 33 wt% silica, the 1st cycle with a displacement of 2.5 mm is recoverable with a maximum load of 0.11 N (after subtracting the base initial load of 0.14 N), while the 2nd cycle with a total displacement of 5 mm is beyond elastic response with a residual displacement of 1.4 mm after complete unloading (Fig. 4a). The concept of mechanical recoverability derives from the microstructural resilience in the fibrous composite

membrane, which is affected by silica aerogel concentrations and hierarchical architectures. The load-displacement cycles for the nanocomposite membranes with 33 wt% silica aerogels (0.3 mm, 0.6 mm, and 0.8 mm thickness) are compared in Fig. 4b, revealing that a thicker sample bears higher load forces during the loading-unloading cycle.

Membranes with thickness below 0.8 mm maintain their recoverability, while membranes with thickness in the region of 0.8 mm have a residual displacement of 1.3 mm. The loss of recoverability for thicker samples could be the result of stress concentration among the non-homogeneous microstructures. The flexibility performance could be quantified by the parameters of bending rigidity and active collateral rate calculated from the load-displacement cycle of the membranes.⁴ For membranes with thickness in the region of 0.3 mm and different concentrations of silica aerogel materials (33 wt%, 40 wt%, and 67 wt%), the bending rigidity of the membrane with 40 wt% silica aerogels shows a higher value of 2.5 cN mm^{-1} compared with 2.1 cN mm^{-1} for the membrane



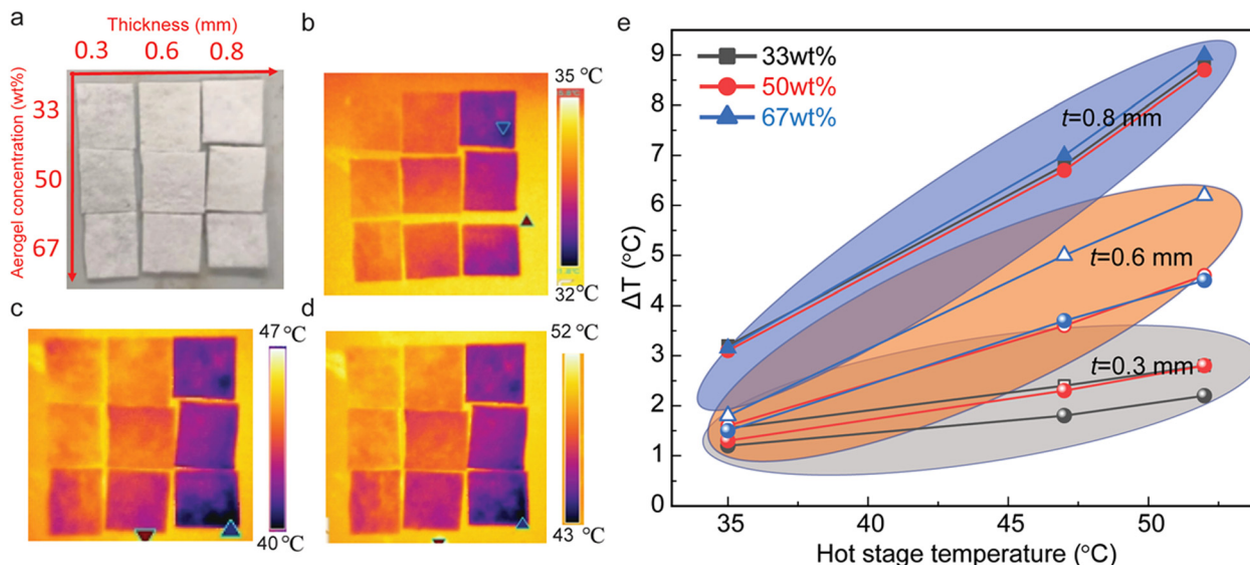


Fig. 3 (a) Optical images of the samples on the hotplate. The scale bar is 10 mm. (b–d) Infrared images of various samples at heating temperatures of 35 °C, 47 °C, and 52 °C. (e) Temperature difference between various composite membranes and the stage vs. hot stage temperature.

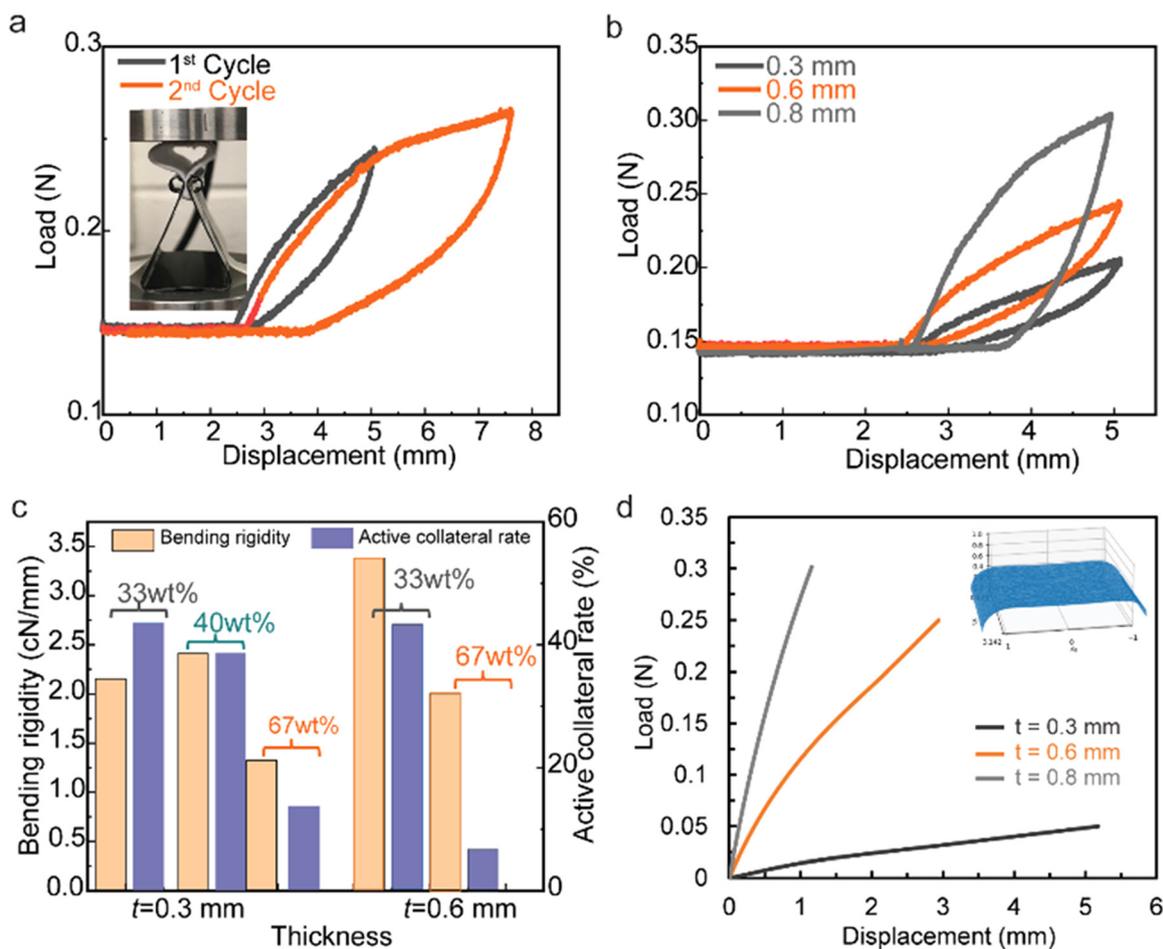


Fig. 4 (a) Two cycle load–displacement curves of 0.6 mm thick membranes with 33 wt% aerogels. (b) Load–displacement curves of 33 wt% aerogel's membrane with thicknesses of 0.3, 0.6, 0.8 mm. (c) Bending rigidity and active collateral rate for membranes at different thicknesses and with different aerogel concentrations. (d) Preliminary load–displacement curves from the modelling of the vertical ring method bending test for the same samples studied in (a–c).



with 33% aerogel, and 1.3 cN mm^{-1} for the membrane with 67 wt% aerogel (Fig. 4c). It should be noted that nanocomposites with superior bending rigidity properties require a synergistic contribution from fiber networks with uniformly distributed silica aerogels. When the thickness increases to 0.6 mm, the bending rigidity increases to 3.4 cN mm^{-1} for the membrane with 33 wt% silica aerogel. The active collateral rate decreases from 43% to 18% when the concentration of silica aerogels increases from 33 wt% to 67 wt%. A smaller active collateral rate indicates larger energy dissipation.

The membrane bending of the material in the vertical ring experiments is modeled using finite element methods. Simulations were conducted with a nonlinear thin-shell assumption and *via* an enriched mixed-element formulation implemented in the finite element library FEniCS.^{23–25} Following this,²⁶ cubic bubble shape functions and reduced integration are employed to resolve the numerical locking issue in this simulation. The top surface of the simulated membrane is subjected to a uniformly distributed compression load. The bottom edges are modeled as a roller support with zero displacement boundary conditions in the vertical direction and allowing it to slide horizontally. To model the increase in the contact area between the plate and the membrane's surface during the vertical ring experiment, the applied load is increased linearly with time. Fig. 4d shows the load-displacement computed from the finite element simulations for three different sample thicknesses. The deformed configuration of the midplane of the top half of the membrane in this figure agrees with the deformed samples in the experiment shown in Fig. 4a. Both the experimental and numerical results demonstrate a nonlinear increase in membrane stiffness with sample thickness.

4 Conclusions

In conclusion, flexible, hydrophobic, and thermal-insulating composite membranes consisting of ceramic-based fibers and silica aerogels featuring a three-dimensional-type network structure are reported.

Moreover, through effective control of the homogeneous microstructure across the membrane nanocomposite, as a result of the combination of short fibers and aerogel precursors under meticulously optimized processing parameters, a low thermal conductivity of $11.4 \text{ mW m}^{-1} \text{ K}^{-1}$, hydrophobicity with a water contact angle of 144° , and mechanical flexibility with a bending rigidity of 1.25 cN mm^{-1} are successfully accomplished.

Fabricated through a scalable, vacuum-filtration-assisted procedure, the nanofibrous-reinforced silica aerogel composite membranes are potential candidates for advanced thermal management applications.

Author contributions

All authors discussed the results and commented on the manuscript. L. A. and S. R. designed all experiments. L. A. and M. D.

prepared the materials and conducted experiments for investigating their mechanical and thermal properties, as well as structural and physicochemical characterization. J. T. and D. F. conducted and supervised the modelling of the mechanical property study, respectively. S. R. conceived the project and directed the research. All authors contributed to the writing and editing of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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